Compound	R	R <sup>1</sup>	S 634- 33	$X^a$	Retention time, min <sup>b</sup>		
			Sequence, 5' to 3'		Gradient 1	Gradient 2	
8	TMT	н	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	0	32.8	_	
9	TMT	Н	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	27.7	23.8	
10	TMT	Pyr	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	-	30.2, 30.7	
11	ТМТ	Pyr	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	О	-	29.1	
12	TMT	Flu	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	33.4, 34.2	_	
13	ТМТ	Flu	AGCT <sub>2</sub> C T <sub>3</sub> GCACA TGTA <sub>3</sub> <sup>c</sup>	s	35.9, 36.3	_	
			<seq. 3="" id.="" no.=""></seq.>				
14	TMT	Flu	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	О	32.9	_	
15	TMT	$C_2H_5$	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	О	33.0	_	
16	Н	Н	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	0	18.0	_	
17	Н	Н	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	S	19.6	16.4	
18	Н	Pyr	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	_	25.0	
19	Н	Pyr	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	О	_	24.5	
20	Н	Flu	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	23.4	-	
21	Н	Flu	AGCT₂C T₃GCACA TGTA₃ <sup>c</sup>	S	28.5	_	
			<seq. 3="" id.="" no.=""></seq.>				
22	Н	Flu	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	О	22.7	-	
23	н	C <sub>2</sub> H <sub>5</sub>	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	o	21.5	_	
24	_	Н	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	o	16.2	_	
25	-	н	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	18.0	15.2	
26	-	Pyr	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	22.6	17.1	
27	-	Pyr	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	o	22.0	16.7	
28	-	Flu	ATGCAT <sub>2</sub> CTGC <sub>5</sub> A <sub>2</sub> G <sub>2</sub> A <seq. 2="" id.="" no.=""></seq.>	s	19.1	14.5	
29	_	Flu	AGCT <sub>2</sub> C T <sub>3</sub> GCACA TGTA <sub>3</sub> <sup>c</sup>	s	21.7	_	
			<seq. 3="" id.="" no.=""></seq.>				
30	-	Flu	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	o	18.1	-	
31	-	C <sub>2</sub> H <sub>5</sub>	T <sub>12</sub> <seq. 1="" id.="" no.=""></seq.>	o	17.9	_	

<sup>a</sup> All oligonucleotides contained uniform, either phosphate (X=O) or phosphorothioate (X=S) backbone; <sup>b</sup> For HPLC conditions, consult Experimental Procedures; <sup>c</sup> 2'-O-(2-methoxyethyl) ribonucleotide residues are italicized; C stands for 5-methyl-2'-O-(2-methoxyethyl)cytidine residue.

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Please delete the paragraph spanning page 49 of the specification and replace it with the following new paragraph.

Table 3.ESMS data oligonucleotides 33-40.<sup>a</sup>

	Sequence, 5' to 3'	Backbone	ESMS, found	Molecular Formula	ESMS, calculated
33	DMTr-TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpO	P=O	6365.2	$C_{211}H_{263}N_{71}O_{122}P_{20}$	6365.2
	<seq. 4="" id.="" no.=""></seq.>				
34	DMTr-TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpS	P=O	6381.7	$C_{211}H_{263}N_{71}O_{121}P_{20}S$	6381.3
	<seq. 5="" id.="" no.=""></seq.>		0301.7	C2[[11263147] O12[1 203	0301.3
35	DMTr-TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpS	P=S	6686.9	$C_{211}H_{263}N_{71}O_{102}P_{20}S_{20}$	6686.5
	<seq. 5="" id.="" no.=""></seq.>				
36	TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpO	P=O	6062.2	$C_{190}H_{245}N_{71}O_{120}P_{20}$	6062.8
	<seq. 6="" id.="" no.=""></seq.>				
37	TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpS	P=O	6079.4	$C_{190}H_{245}N_{71}O_{119}P_{20}S$	6078.9
	<seq. 7="" id.="" no.=""></seq.>				
38	TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpS	P=S	6384.7	$C_{190}H_{245}N_{71}O_{100}P_{20}S_{20}$	6384.2
	<seq. 7="" id.="" no.=""></seq.>				
39	TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpS-Pyr	P=O	6350.9	$C_{209}H_{258}N_{72}O_{120}P_{20}S$	6350.2
	<seq. 8="" id.="" no.=""></seq.>				
40	TGCATC <sub>5</sub> AG <sub>2</sub> C <sub>2</sub> AC <sub>2</sub> ATpS-Flu	P=O	6467.0	$C_{212}H_{258}N_{72}O_{125}P_{20}S$	6466.2
	<seq. 9="" id.="" no.=""></seq.>				

Please delete the paragraph spanning page 77 of the specification and replace it with the following new paragraph.

Table 9: Cholesterol-Conjugated Oligonucleotides (4)

Cmpd	Sequence (5' – 3')	Backbone	Quantity	MW		More
				calc	Found	Info
				•		